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- (54) Organic light emitting diode device with three component emitting layer
- (57) An organic light emitting device includes a substrate, an anode and a cathode disposed over the substrate; a luminescent layer disposed between the anode and the cathode wherein the luminescent layer includes a host and at least one dopant; the host of the luminescent layer being selected to include a solid organic material comprision a mixture of at least two components.

wherein the first component of the mixture is an organic compound that is capable of transporting both electrons and holes and that is substantially non-polar, and the second component of the mixture is an organic compound that is more polar than the first component; and the depart of the luminescent layer being selected to produce light from the light emitting device.

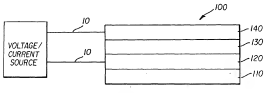


FIG I

Description

[0001] The present invention relates to organic light emitting diode devices and more particularly to the design of the composition of the organic layers for improvements in luminance efficiency.

[0002] Organic light emitting diodes (OLED), also known as organic electroluminescent (EL) devices, are a class of electronic devices that emit light in response to an electrical current applied to the device. The structure of an OLED device generally includes an anode, an organic EL medium, and a cathode. The term, organic EL medium, herein refers to organic materials or layers of organic materials disposed between the anode and the cathode in the OLED device. The organic EL medium may include low molecular weight compounds, high molecular weight polymers, oligimers of low molecular weight compounds, or biomaterials, in the form of a thin film or a bulk solid. The medium can be amorphous or crystalline. Organic electroluminescent media of various structures have been described in the prior art, Dresner, in RCA Review, 30, 322 (1969), described a medium comprising a single layer of anthracene film, Tang and others, in Applied Physics Letters, 51, 913 (1987), Journal of Applied Physics, 65, 3610 (1989), and commonlyassigned US-A-4,769,292, reported an EL medium with a multi-layer structure of organic thin films, and demonstrated highly efficient OLED devices using such a medium. In some OLED device structures the multi-layer EL medium includes a hole transport layer adjacent to the anode, an electron transport layer adjacent to the cathode, and disposed in between these two layers, a luminescent layer. Furthermore, in some preferred device structures, the luminescent layer is constructed of a doped organic film comprising an organic material as the host and a small concentration of a fluorescent compound as the dopant. Improvements in EL efficiency and chromaticity have been obtained in these doped OLED devices by selecting an appropriate dopant-host composition. Often, the dopant, being the dominant emissive center, is selected to produce the desirable EL colors. Examples of the doped luminescent layer reported by Tang and others in commonly-assigned US-A-4,769,292 and by Chen and others in commonly-assigned US-A-5.908.581 are: tris(8-quinolinol)aluminum (AIQ) host doped with coumarin dves for green emitting OLEDs; and AIQ doped with 4-dicyanomethylene-4H-pyrans (DCMs) for orange-red emitting OLEDs. In addition to enhancements in color and luminance efficiency, there are other significant benefits in using a doped luminescent layer. Shi and others, in commonly-assigned US-A-5.593,788, disclosed that the use of a doped luminescent layer significantly improved the stability of OLED devices. In Shi's disclosure, a long operational life was obtained in an OLED device by using a quinacridone compound as the dopant in an AIQ host. The quinacridone dopant produced a green emission with a high luminance efficiency. Bryan and others in commonly-assigned US-A-5,141,671, disclosed a luminescent layer containing perylene or a perylene derivative as a dopant in a blue emitting host. They showed that a blue emitting OLED device with an improved blue hue and an improved operational stability was obtained. In both disclosures, the incorporation of selected fluorescent dopants in the luminescent layer is found to improve substantially the overall OLED device performance parameters.

[0003] The most common formulation of the doped luminescent layer includes only a single dopent in a host matrix.
However, in a few instances, incorporation of more than one obpant in the luminescent layer was found to be beneficial in improving the flue. One such instance was reported by Hamada and others in Applied Phys. Lett. 75, 1882 (1999).
Using a luminescent layer containing rubrene, a yellow entiting dopant, and DCJ 4-(doyanomethylene)2-mothyl6/2(4-(s)licity)ethentyl-41-Pyran, a red emitting dopant, in an AIQ most, Hamada and others were able to produce a demitting OLED device with excellent chromaticity. Furthermore, the red hus from the DCJ dopant remains essentially
unchanged regardless of the intensity of light output from the OLED. In contrast, with only DCJ dopant in the AIQ host,
the color of the light emitted from the dopant DCJ was noticeably blue-shifted, producing a less than desirable orange,
rather than red, hue. With the dual obpant system, rubrene functions as a co-dopant in mediating energy transfer from
the AIQ nost to the DCJ emitter. In either single dopant or dual obpant systems, it has been noted that the luminance
ficiency tends to decrease with increasing brightness, that is with increasing current density. With red-emitting dopants, the hue usually shifts towards orange with increasing current density.

[0004] Although EL efficiency, color, and stability have been improved significantly using doped luminescent layers of various compositions, the problem of decreasing EL efficiency with increasing light output or drive current density persists, particularly in red-rentiting OLED devices.

[0005] It is an object of the present invention to provide OLED devices with improved luminance efficiency that is essentially independent of the intensity of the light output from the OLED.

[0006] It is another object of the present invention to provide specifically red OLED devices with an improved luminance efficiency that is essentially independent of the intensity of the light output.

[0007] It is a further object of the present invention to provide specifically red OLED devices with chromaticity essentially independent of the light output.

[0008] These objects are achieved in an organic light emitting device comprising:

- (a) a substrate
- (b) an anode and a cathode disposed over the substrate;

(c) a luminescent layer disposed between the anode and the cathode wherein the luminescent layer includes a host and at least one dopant:

(d) the host of the luminescent layer being selected to include a solid organic material comprising a mixture of at least two components wherein;

(i) the first component of the mixture is an organic compound that is capable of transporting both electrons and holes and that is substantially non-polar; and

(ii) the second component of the mixture is an organic compound that is more polar than the first component;

and

(e) the dopant of the luminescent layer being selected to produce light from the light emitting device,

[0009] An advantage of the present invention is that, with an appropriate selection of the first and second host com-

ponents and the dopants in the luminescent layer, OLED devices with high luminance efficiencies are produced. [0010] Another advantage of the present invention is that it provides OLED devices with a luminance efficiency which is relatively independent of the intensity of the light output from the OLED.

[0011] Another advantage of the present invention is that it provides an OLED device with a chromaticity essentially independent of the intensity of the light output from the OLED

[0012] Another advantage of the present invention is that it provides a red emitting OLED with excellent efficiency

and chromaticity. [0013] FIG. 1 is schematic structure of an OLED with an organic EL medium; and

[0014] FIG. 2 and FIG. 3 are two schematic OLED structures showing two different configurations of the organic EL

[0015] The drawings are necessarily of a schematic nature, since the individual layers are too thin and the thickness differences of the various elements too great to permit depiction to scale or to permit convenient proportionate scaling. [0016] FIG. 1 illustrates the structure of an OLED device of the simplest construction practiced in the present invention. In this structure, OLED device 100 includes an anode 120, an EL medium 130, and a cathode 140, disposed upon a substrate 110. In operation, an electrical current is passed through the OLED by connecting an external current or voltage source with electrical conductors 10 to the anode and the cathode, causing light to be emitted from the EL medium. The light may exit through either the anode or the cathode or both as desired and depending on their optical transparencies. The EL medium includes a single layer or a multi-layer of organic materials.

[0017] FIG. 2 rates the structure of another OLED device of the present ion. In this structure, OLED device 200 includes a substrate 210 and an EL medium 230, disposed between anode 220 and cathode 240. EL medium 230 includes a hole-transport layer 231 adjacent to the anode, an electron transport layer 233 adjacent to the cathode, and a luminescent layer 232 disposed between the hole-transport layer and the electron-transport layer. In operation, an electrical current is passed through the OLED device by connecting an external current or voltage source with electrical conductors 10 to the anode and the cathode. This electrical current, passing through the EL medium, causes light to be emitted primarily from the luminescent layer 232. Hole-transport layer 231 carries the holes, that is, positive electronic charge carriers, from the anode to the luminescent layer. Electron-transport layer 233 carries the electrons, that 40 is, negative electronic charge carriers, from the cathode to the luminescent layer 232. The recombination of holes and

electrons produces light emission, that is, electroluminescence, from the luminescent layer 232. [0018] FIG. 3 illustrates yet another structure of an OLED device of the present invention. In this structure, OLED device 300 includes a substrate 310 and an EL medium 330 disposed between anode 320 and cathode 340. EL medium 330 includes a hole injection layer 331, a hole transport layer 332, a luminescent layer 333, an electron transport layer 334, and an electron injection layer 335. Similarly to OLED device 200 of FIG. 2, the recombination of electrons and holes produces emission from the luminescent layer 333. The provision of the hole-injection layer 331 and the electron injection layer 335 serves to reduce the barriers for carrier injection from the respective electrodes. Consequently, the drive voltage required for the OLED device can be reduced.

[0019] According to the present invention, the luminescent layer (either layer 232 of FIG. 2 or layer 333 of FIG. 3) is primarily responsible for the electroluminescence emitted from the OLED device. One of the most commonly used formulations for this luminescent layer is an organic thin film including a host and one or more dopants. The host serves as the solid medium or matrix for the transport and recombination of charge carriers injected from the anode and the cathode. The dopant, usually homogeneously distributed in the host in small quantity, provides the emission centers where light is generated. Following the teaching of the prior art, the present invention uses a luminescent layer including a host and a dopant, but it distinguishes over the prior art that the host of the present invention is a mixture having at least two components, each component having specific electronic properties. The selection of these host components and compatible dopant materials is in accordance with the following criteria:

- 1. The host is a solid organic thin film having at least two uniformly mixed host components:
- The first host component is an organic compound capable of transporting both electrons and holes, and the molecular structure of this organic compound is substantially non-polar;
- The second host component is an organic compound having a molecular structure more polar than that of the first host component.
 - 4. The dopant is an organic luminescent compound capable of accepting the energy released from the recombination of electrons and holes in either the first or second host components, and emitting the energy as light.
- [0020] Following the selection orther in of this invention, OLEO devices have been constructed having excellent luminance efficiency and chromaticity. More importantly, the luminance efficiency measured in candetes per amper is essentially constant over a large range of brightness or current densities. This is a distinct advantage over the prior art, where the luminance efficiency often decreases, or otherwise varies, with increasing brightness or current density. Another important advantage is that the chromaticity also remains essentially constant, independent of the brightness or current density.
- 15 [0021] Preferred materials for the first host component of the luminoscent layer of this invention include a class of compounds which, for the purpose of this invention, will be referred to as between old compounds. The bearzand/compounds comprise polycyclic hydrocarbons (PAH) and combinations of two or more PAH. In benzenoid compounds formed by combination of two or more PAH the PAH are connected together via a single chemical bond or linked via a saturated or unsaturated hydrocarbon group. The list of PAH useful as building blocks for benzenoid compounds includes.
 - 1. Benzene
 - 2. Naphthalene
- 3. Anthracene
 - Phenanthrene
 - Tetracene
 - Pyrene
 Pentacene
 - 8. Perylene
 - 9. Coronene

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- 10. Chrysene
- 11. Picene
- 12. Pericyclene
- 35 [0022] Any of the above listed PAH, and any benzenoid compounds formed by the combination of one or more of the above listed PAH, are useful as the first host component, provided that the compounds are also film-forming materials at room temperature. Benzenoid compounds containing a total of 5 or more six-membered rings in PAH groups are usually film-forming
- [0023] Representative examples of useful benzenoid compounds in which the PAH are connected together via single demical bonds include:
 - 1) Hexaphenylbenzene
 - Benzene-anthracene-benzene (for example 9.10-diphenylanthracene)
 - 3) Naphthalene-anthracene- naphthalene. (for example 9.10- bis(2-naphthyl)anthracene)
 - 4) (Benzene)2-tetracene-(benzene)2 (for example rubrene)
 - 5) Anthracene-anthracene (for example, bianthryl)
 - 6) Anthracene-anthracene-anthracene (for example, 9.10-dianthrylanthracene)
 - 7) Pyrene-perylene
 - 8) Perylene-anthracene
 - In the above examples, a hyphen represents a single chemical bond between PAH moieties. Useful benzenoid compounds include compounds including PAH groups linked by one or more hydrocarbon groups. Any of the above benzenoid compounds substituted with one or more hydrocarbon groups are useful.
- [0024] Particularly preferred materials for the first host component of the luminescent layer of this invention include 55 benzenoid compounds of the following structure:

 R^1 R^2

wherein:

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substituents R¹, R², R³ and R⁴ are each individually hydrogen, or alkyl of from 1 to 24 carbon atoms, or aryl, or substituted aryl of from 5 to 30 carbon atoms.

[0025] The chemical names and their abbreviations of specific examples of the particularly preferred materials in-

9,10-Diphenylanthracene (DPA)

9,10-Bis(2-naphthalenyl)anthracene (ADN)

2-(1,1-dimethylethyl)-9,10-bis(2-naphthalenyl)anthracenc (TBADN)

9,10-Bis[4-(2,2-diphenylethenyl)phenyl]anthracene

[0026] One particular selection criterion for the first nest component is that the organic compound should have a molecular structure that is substantially non-polar. The polarity of a compound is a measure of the distribution of electrical charge in the molecule. A non-polar or substantially non-polar molecule has a structure wherein every atom is approximately electrically neutral. In contrast, a polar compound has a moleculer structure wherein some atoms have a partial positive or negative charge. The polarity of a notecule can, in some instances, be indicated by the magnitude of the cipole moment, defined in units of Debye. For reference, the values of the cipole moments of a large number of organic compounds can be found in the Handbook of Chemistry and Physics, 51st edition, page E70. A non-polar molecule usually has a dipole moment less than 1.0 Debye.

[0027] For some organic compounds, the molecular structure is such that the average dipole moment of the molecular can be relatively small when though the molecular structure may contain various moleties or group of atoms that are rather polar as individual groups. The reason is that the gipole moments of the individual more molecular structure may oppose each other, resulting in cancellation of these dipole moments. The values of the dipole moment of various groups of atoms within a molecule, known as group dipole moments, can be found in the book Dipole Moments in Organic Chemistry by V. I. Minkin, O. A. Osipov, and Y. A. Zhdanov. Further examples of the use of net and group dipole moments to describe the nonpolar or polar character of molecules useful in OLED are available in Young and Floragradi. Journal of Playsical Chemistry, 99, 4200 (1995).

[0028] As used herein, the dipole moment of a molety or a group of atoms within a molecule is referred to as the group dipole moment. The dipole moment of the entire molecule is termed the net dipole moment of the molecule. For an organic compound to be substantially non-polar as used herein, the group dipole moments or the net dipole moment of a molecule should be less than one Debye.

[0028] Almost all benzenoid compounds found useful as the first host component in the present invention have a dipole moment of less than one Debye, and groups of almos within these compounds also have dipole moments of less than one Debye. Many representative benzenoids, such as naphthalene, anthrezene, and perylene, have practically zero group and ned dipole moments. Other organic compounds meeting such a non-poleur critical are usual as well.

[0030] Materials for the second host component of the luminescent layer of the present invention include organic compounds that have a dipole moment at its higher than 4 of the first host component. As defined previously, the values of the adoption of the present invention include organic compounds that have passed and the group dipole moment of molecules can be quite different. For an organic compound to be useful as the second host component, it is only necessary that any individual group dipole moment of the molecule be greater than any individual group alpice moment or the dipole moment of the molecule used for the first host component. The value of 0.5 Debye or larger is preferred for the individual group or a dipole moment for molecule used for the first host component. The value of 0.5 Debye or larger is preferred for the individual group or a dipole moment for molecule useful as the second host component in the luminescent layer.

[0031] Preferably the second host component should have a bandgap that is less than that of the first host component. The bandgap is defined as the energy needed to bring an electron from the highest occupied molecular orbital to the lowest unoccupied molecular orbital of the molecule. This condition ensures that energy transfer from the first host component to the second host component is favorable and the energy resulting from the recombination of electrons and holes in the first host component can be transferred to the second host component and subsequently to the light producing dopants.

[0032] Another reason for selecting a material of lower bandgap for the second host component is that it may also serve as a hole trap, an electron trap, or both. Trapping injected carriers directly in the second host component is beneficial as it promotes electron-hole recombination in this component, shortcutting the need for carrier recombination in the first host component is needed for carrier transport only.

[0033] Materiais useful as the second host component in the luminescent layer include benzenoids that are rendered more polar with one or more electron-dending or electron-withdrawing moieties in the molecular structure. Electron-donating moieties include: armino, alkylamino, anylamino, diskylamino, disrylamino, methoxy, and phenoxy. Electron-withdrawing moieties include: organo, nitro, flutor-or, tohro, kelo, cachovy, and pvindy.

[0034] Another class of materials useful as the second host component includes benzenoids that contain heterocyclic

structures. These structures include benzoxazolyl, and thio and amino analogs of benzoxazolyl of following general molecular structure:

wherein

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Z is O, NR" or S;

R and R', are individually hydrogen; alkyl of from 1 to 24 carbon atoms; anyl or hetero-atom substituted anyl of from 5 to 20 carbon atoms; halo, or atoms necessary to complete a fused aromatic ring; and R' is hydrogen; alkyl of from 1 to 24 carbon atoms; or anyl of from 5 to 20 carbon atoms; or anyl of from 5 to 20 carbon atoms; or anyl of from 5 to 20 carbon atoms.

[0035] The dipole moment for the benzimidazole group of this structure is estimated to be approximately 4 Debyes from peopled lightcarture [Tables of Experimental Dipole Moments, detledy A. L. McCiellan, Fereman, 1983, Page 238]. [0036] An experimental people of the properties of the second host component is the oxinoid compounds. Exemplary of contembeted professional compounds are those satisfying the following structural formula:

$$\left[\begin{array}{ccc} \overset{z}{\swarrow} & & \\ & \overset{z}{\searrow} & \\ & & & \end{array}\right]_n \quad \longleftrightarrow \quad \left[\begin{array}{ccc} \overset{z}{\swarrow} & & \\ & \overset{z}{\searrow} & \\ & & & \end{array}\right]_{k^{+n}}$$

wherein

Me represents a metal;

n is an integer of from 1 to 3; and

Z independently in each occurrence represents the atoms completing a nucleus having at least two fused aromatic rings.

[0037] From the foregoing it is apparent that the metal can be monovalent, divalent, or trivalent metal. The metal can, for example, be an alkalif metal, such as ithium, sodium, or potassium; an alkaline earth metal, such as magnesium or calcium; or an earth metal, such as boron or aluminum. Generally any monovalent, divalent, or trivalent metal known to be a useful chelating metal can be employed.

[0038] Z completes a heterocyclic nucleus containing at least two fused aromatic rings, at least one of which is an azole or azine ring. Additional rings, including both aliphatic and aromatic rings, can be fused with the two required rings, if required. To avoid adding molecular bulk without improving on function the number of ring atoms is preferably maintained at 18 or less.

[0039] Illustrative of useful chelated oxinoid compounds and their abbreviated names are the following:

Tris(8-quinolinol)aluminum (AIQ) Bis(8-quinolinol)-magnesium (MgQ) Tris(8-quinolinol)indium (InQ) 8-quinolinol lithium (LiQ)

[0040] The material selection criteria for the dopant in the luminescent layer are: 1) the dopant molecule has a high efficiency of fluorescence or phosphorescence in the luminescent layer, and 2) it has a bandgap smaller than that of the both the first and second host materials.

[0041] For red emitting OLEDs, a preferred class of dopants of this invention has the general formula:

15 wherein:

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R1, R4, R3, and R1 are individually allyl of from 1 to 10 carbon atoms;

R5 is alkyl of from 2 to 20 carbon atoms, anyl, storically hindered anyl, or heteroanyl:
and R5 is alkyl of from 1 to 10 carbon atoms, or a 5- or 6-membered carbocyclic trig connecting with R5.

[0042] These materials possess fluorescence efficiencies as high as unity in solutions and emit in the orange-red spectral region. Appresentative materials of this closes and their abbreviated names include:

N°C OH DCM

NC CN DCJ

NC_CN . DCJT

[0043] Another class of fluorescent materials that are useful as the dopant in the present invention includes compounds of the formula:

wherein:

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X = S, or O

R₁ and R₂ are individually alkyl of from 1 to 20 carbon atoms, anyl or carbocyclic systems;

 R_3 and R_4 are individually alkyl of from 1 to 10 carbon atoms, or a branched or unbranched 5 or 6 member substituent ring connecting with R_1 , R_2 respectively; and

Rs and Rs are individually alkyl of from 1 to 20 carbon atoms, which are branched or unbranched.

[0044] Representative materials of this class and their abbreviated names include:

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- 49 [0045] The composition of the luminescent layer of this invention is such that the first host component constitutes the largest volume fraction, followed by the second host component. The depart constitutes the smallest volume fraction. The preferred range for the first host component is no more than 95% but no less than about 50% by volume. The preferred concentration range for the second host component is no more than about 40% but no less than about 5% by volume. The preferred concentration range for the depart is between 0.1% to 10% by volume. The thickness is the second that the preferred range is the thing the second post component is not more than about 40% but no less than about 5% by volume. The thickness is the second that the preferred range is the thing the preferred range is the them. The thickness is this range is sufficiently large to enable recombination of charge carriers and, therefore, electroluminescence to take place axious/volume to the preferred range is between 100 Angstroms and 500 Angstroms, where the overall OLED device performance parameters, including drive voltage, are columnal.
 - [0046] A useful method for forming the luminescent layer of the present invention is by vapor deposition in a vacuum chamber. This method is particularly useful for fabricating oLLED devices, where the layer structure, including the organic layers, can be sequentially deposited on a substate without significant interference among the layers. The thickness of each individual layer and its composition can be precisely controlled in the deposition process. To produce the desired composition of the luminescent layer, the rate of deposition for each component is independently controlled using a deposition rate monitor.
- 55 [0047] Returning to FiG. 2, hole-transport layer 231 and electron-transport layer 233 provide the functions of transporting holes and electrons, respectively, to the luminescent layer 232. The use of these layers and their material compositions in OLED devices have been disclosed by Tang et all in commonly-assigned US-A-4.769,292, include herein by reference. A fycical hole-transport layer includes the hole-transporting compound NPB, NN-bisf1-nachthyl)

-N.N'-diphenvlbenzidine.

[0048] Returning to FIG. 3, hole-injection layer 331 and electron-injection layer 335 provide the functions of improving the hole injection from the anode and electron injection from the cathode 340, respectively. The use of a hole-injection layer in OLED devices has been disclosed by Van Slyke and others in commonly-assigned US-A-4,720,432, included herein by reference. The use of an electron-injection layer has been disclosed by Hung et al. in commonly-assigned US-A-5,776,52, also included herein by reference.

Comparative Example 1

[0049] An OLED device was prepared as follows. A glass substrate coated with a transparent indium-tin-oxide (ITO) conductive layer was cleaned and dried using a commercial glass scrubber fool. The ITO surface was subsequently treated with an oxidative plasma to condition the surface as an anode. The following layers were deposited in the following sequence by sublimation from a heated crucible boat in a conventional vacuum deposition chamber under a vacuum of approximately 10⁴ forr. (1) a hole-transport layer, 700Å thick, consisting of NPB, (2) a luminescent layer, 350 Å thick, consisting of TBADN as the sole host material and 1½ DCJTB as the dopant, (3) an electron-transport layer, 400 Å thick, consisting of AIQ, and (4) a cathode, approximately 2200 Å thick, consisting of an alloy of magnesium and silver with a MOAA volume ratio of about 10:1.

[0050] The EL characteristics of this device were evaluated using a constant current source and a photometer. The efficiency, CIE coordinates, and drive voltage at a reliatively low current density, 1 mA/cm², and a relatively high current density, 10 mA/cm², were measured. Their values are shown in Table I.

Example 2

[0051] An CLED device similar to that of Comparative Example 1 was constructed, except that in the luminescent layer. TRADN is the material for the first host component, All or the material for the second host component, and or the material for the second host component, and or CLTB on a volume basis are in the ratio SCLTB is the fluorescent dopant. The relative amounts of TRADN, AlO, and DCJTB on a volume basis are in the ratio SR101. The CL characteristics of this device are also shown in Table.

Examples 3 - 5

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[0052] OLED devices similar to that of Example 2 were constructed, except that the relative amounts of TBADN, AIQ, and DCJTB in the luminescent layer are in the ratio 74:25:1 for Example 3, 49:50:1 for Example 4, and 25:74:1 for Example 5. The EL characteristics of these devices are also shown in Table I.

35 Comparative Example 6

[0053] An OLED device similar to that of Comparative Example 1 was constructed, except that the sole host material in the luminescent layer was AlQ rather than TBADN. The EL characteristics of this device are also shown in Table I.

Table I. Compositions and EL properties of OLED

					_		_		_		_			
Drive	voltage	B	4.1	9.6	5.6	11.2	5.6	11.3	5.7	11.5	6.4	12.2	6.9	12.3
CIEy			0.448	0.457	0.392	0.399	0.381	0.388	0.368	0.376	0.356	0.366	0.350	0.368
CIEX			0.534	0.521	0.603	0.595	0.615	909.0	0.629	0.620	0.642	0.629	0.648	0.628
Efficiency	(cd/A)		3.54	3.85	5.93	4.98	5.75	4.62	4.73	3.53	3.35	2.45	2.49	1.60
Current	density	(mA/cm²)	-	100	-	100	-	001	-	001	-	100	-	100
Dopant	DCJTB	%loA		-	-	-	-	-	-	-		-	-	-
Second host	AIQ	%lo/		>	5	2	30	3	9	3	ř	ţ	96	66
First host	TBADN	%loA	8	3	8	8	77	ţ	Ç	î	3.6	3		>
Example			-	-	,	7	,	0		+	,	,	9	0
	First host Second host Dopant Current Efficiency CIEx CIEy	First host Second host Dopant Current Efficiency CIEX CIEY TBADN AlQ DCJTB density (cd/A)	First host Second host Dopant Current Efficiency CIEx CIEy TBADN AlQ DCJTB density (cd/A) Cd/A) Vol% Vol% Vol% (mA/cm²) CEVA	First host Second host Dopant Current Efficiency CIEx CIEy TBADN AlQ DCJTB density (cd/A) Cd/A) Vol% Vol% Vol% (mA/cm²) 1 3.54 0.534 0.448	First host Second host Dopant Current Efficiency CIEX CIEY TBADN AlQ DCJTB density (cd/A) Vol9% Vol9% (mA/cm²) 99 0 1 1 3.54 0.534 0.448	First host Second host Dopant Current Efficiency CIEx CIEy TBADN AlQ DCJTB density (cd/A) Vol96 Vol96 Vol96 (mA/cm²) 99 0 1 1 3.54 0.534 0.448 99 0 1 1 0.659 0.651 0.457	First host Second host Dopant Current Efficiency CLEX CLEY TBADN AlQ DCITB density (cd/A) Vol% Vol% (mA/cm²) 3.54 0.534 0.448 99 0 1 1 5.93 0.603 0.392 89 10 1 100 4.98 0.595 0.399	First host Second host Dopant Current Efficiency CIEX CIEY TBADN AlQ DCITB density (cd/A) Vol% Vol% (mA/cur) 3.54 0.534 0.448 99 0 1 100 3.85 0.521 0.4457 89 10 1 5.93 0.603 0.392 89 10 1 100 4.98 0.595 0.399 74 75 75 75 0.615 0.381 75 75 75 0.615 0.615 75 75 75 0.615 0.615 75 75 75 0.615 0.615 75 75 75 0.615 0.615 75 75 0.615 75 75 0.615 0.615 75 75 0.615 0.615 75 75 0.615 0.615 75 75 0.615 0.615 75 75 0.61	First host Second host Dopant Current Efficiency CIEX CIEY TBADN Alto DCITB density (cd/A) Vol% Vol% (mA/cur) 3.54 0.534 0.448 99	First host Second host Dopant Current Efficiency CIEX CIEY TBADN AlQ DCITB density (cd/A) Voll% Voll% (mA/cur) 99	First host Second host Dopant Current Efficiency CIEX CIEY TBADN AlQ DCITB density (cd/A) Volv6 Volv6 (mA/cmr) 99	First host Second host Dopant Current Efficiency CIEX CIEY	First host Second host Dopant Current Efficiency CIEX CIEY	First host Second host Dopant Current Efficiency CIEX CIEY

Comparative Examples I and 6 show that the luminance efficiency of an OLED device using a single host material is reliatively low: less than 4 cd/A for Example 1 where the sole host materials is TBADN, and less than 2 5 cd/A for Example 55

55 All the sole host materials is A/D. In contrast, the luminance efficiency of an OLED device with a mixture of first and second host materials is significantly higher, above 3 cd/A for all examples and as high as 5.93 cd/A for Example 2. In Examples 2 - 5, the first host material, TBADN, is a substantially nonpolar benzenoid hydrocarbon. The second host material A/Q, is relatively polar, with a dipole moment of 5.5 Debye. The bandgap values for TBADN, A/Q, and

DCJTB are respectively, 3.2, 2.8, and 2.4 electron volts

[0054] The luminance efficiency at both low and high current densities is generally the highest in OLED devices with a mixture of host components in the luminescent layer [Examples 2, 3, 4]. For OLED devices with a single host component [Comparative Examples 1 and 6], the luminance efficiencies are considerably lower. Moreover, the OLED device of Example 6 exhibits a major drop in luminance efficiency with increasing current density. In addition, according to the color coordinates CIEx and CIEy, the color of the OLED devices is red shifted with increasing concentration of the second host component in the luminescent layer.

Examples 7-10

[0055] OLED devices similar to those of Examples 2-5 were constructed, except that the first host material is ADN. Again, the second host material is AIQ, and the fluorescent dopant is DCJTB. The composition of the luminescent layer and the EL characteristics of these devices are shown in Table III. Conclusions similar to those drawn for TBADN:AIQ in Examples 1-6 can be made for the ADN:AIQ mixed host.

Table II. Compositions and EL properties of OLED devices of Examples 7-10

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Drive	voltage (V)		6.9	13.0	7.0	12.8	8.9	12.5	8.9	12.4
CIEy			0.376	0.382	0.371	0.378	0.365	0.374	0.362	0.374
CIEx			0.621	0.614 0.382	0.626	0.618	0.633	0.622	0.635	0.622
Efficiency CIEx	(cd/A)		6.27	6.15	5.78	5.47	5.26	4.41	4.59	3.25
Current	density	(mA/cm²)	L	100	1	100	-	100	_	100
Dopant	DCTTB	%loA	_	•				•	-	
Second host Dopant	AIQ	%loA	10	2	35	3	60	3	7.4	
First host	ADN	Vol%	08	3	2.7	ţ	40	ì	35	3
Example			7		0		0	,	9	2

Comparative Example 11

[0056] An OLED device similar to that of Comparative Example 1 was constructed, except that the sole host material in the luminescent layer was ADN rather than TBADN, and the fluorescent dopant was C-545T rather than DCJTB. The EL characteristics of this device are shown in Table III.

Examples 12-14

[0057] OLED devices similar to that of Comparative Example 11 were constructed, except that the material for the first host component in the luminescent layer is ADN, and the material for the second host component is AIQ. The composition of the luminescent layer and the EL characteristics of these devices are shown in Table III.

Table III Compositions and EL properties of OLED devices of Examples 11-14.

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Drive	voltage (V)		5.0	8.1	4.7	7.9	4.5	8.0	4.2	7.5
CIEy			809.0	0.604	0.634	0.631	0.638	0.636	0.647	0.645
CIEx			0.244	0.241	0.254	0.253	0.255	0.254	0.261	0.259
Efficiency	(cd/A)		2.57	4.83	4.1	6.2	5.4	6.7	6.9	7.3
Current	density	(mA/cm²)	-	100	-	100	-	100	_	100
Dopant	C-545T	Vol%	-		-		-		1	
Second host Dopant	AIQ	%loA	0		5		01		20	
irst host	ADN	%loA	66		94		68		79	
Example			=		12		13		14	

Comparison of Examples 12-14 with Comparative Example 11 shows that the luminance efficiency of an OLED device 55 using a mixture of ADN and A/O as first and second host materials is considerably higher than that using ADN as the sole host material. In addition, the CIEx and CIEy coordinates indicate a more saturated green hue. [00.58] Other features of the invention are included below.

[0059] The organic light emitting device wherein the second component includes a benzenoid substituted with a

donor or an acceptor moiety.

[0060] The organic light emitting device wherein the second component includes organic compounds of molecular structure:

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$$\left[\begin{array}{ccc} \overset{z}{\underset{N_{0}+n}{\longrightarrow}} & & & \\ & \overset{z}{\underset{N_{0}+n}{\longrightarrow}} & & \\ & & & \\ & & & \\ \end{array}\right]_{n}$$

wherein

Me represents a metal:

n is an integer of from 1 to 3; and

Z independently in each occurrence represents the atoms completing a nucleus having at least two fused aromatic rings.

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[0061] The organic light emitting device wherein the second component includes AlQ, InQ. GaQ, or MgQ.

[0062] The organic light emitting device wherein the dopant comprising a compound of the formula:

wherein:

R1, R2, R3, and R4 are individually alkyl of from 1 to 10 carbon atoms;

R5 is alkyl of from 2-20 carbon atoms, aryl, sterically hindered aryl, or heteroaryl;

and R6 is alkyl of from 1 to 10 carbon atoms, or a 5- or 6-membered carbocyclic ring connecting with R5.

[0063] The organic light emitting device wherein the dopant includes DCM, DCJT, or DCJTB. [0064] The organic light emitting device wherein the dopant includes compounds of structure:

wherein:

X = S, or O

R₁ and R₂ are individually alkyl of from 1 to 20 carbon atoms, anyl or carbocyclic systems;

Ra and Ra are individually alkyl of from 1 to 10 carbon atoms, or a branched or unbranched 5 or 6 member substituent ring connecting with R1, R2 respectively; and

R₅ and R₆ are individually alkyl of from 1 to 20 carbon atoms, which are branched or unbranched. [0065] The organic light emitting device wherein the dopant includes C-6, C-545T, or C525T.

Claims

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1. An organic light emitting device comprising:

(a) a substrate:

(b) an anode and a cathode disposed over the substrate;

(c) a luminescent layer disposed between the anode and the cathode wherein the luminescent layer includes a host and at least one dopant;

(d) the host of the luminescent layer being selected to include a solid organic material comprising a mixture

of at least two components wherein;

(i) the first component of the mixture is an organic compound that is capable of transporting both electrons and holes and that is substantially non-polar; and

(ii) the second component of the mixture is an organic compound that is more polar than the first component; and

(e) the dopant of the luminescent layer being selected to produce light from the light emitting device.

The organic light emitting device of claim 1 wherein the second component has a bandgap that is smaller than that of the first component.

The organic light emitting device of Claim 1 wherein the dopant of the luminescent layer has a bandgap that is smaller than those of the first and second host components.

4. The organic light emitting device of claim 1 wherein the dipole moment of the first component is less than 1 Debye.

The organic light emitting device of claim 1 wherein the dipole moment of the second component is greater than 0.5 Debve.

The organic light emitting device of claim 1 wherein the dipole moment of a group of atoms within the second component is greater than 0.5 Debye.

7. The organic light emitting device of claim 1 wherein the dopant is a luminescent organic compound.

8. The organic light emitting device of claim 1 wherein the first host component includes a benzenoid compound.

9. The organic light emitting device of claim 8 wherein the benzenoid compound has a the formula:

wherein

substituents ${\rm R}^1, {\rm R}^2, {\rm R}^3$ and ${\rm R}^4$ are each individually hydrogen, alkyl of from 1 to 24 carbon atoms, aryl, or substituted anyl of from 5 to 30 carbon atoms.

10. The organic light emitting device of claim 1 wherein the first component includes:



9,10-Diphenylanthracene (DPA)

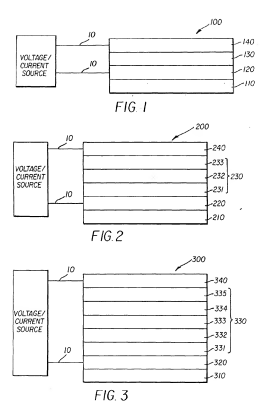


9,10- Bis(2-naphthalenyl)anthracene (ADN)



2-(1,1-dimethylethyl)-9,10-bis(2-naphthalenyl)anthracene (TBADN)

9,10-Bis[4-(2,2-diphenylethenyl)phenyl]anthracene





European Pater Office

EUROPEAN SEARCH REPORT

Application Number EP 01 20 5032

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